

UNITED STATES PATENT AND TRADEMARK OFFICE

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.		
10/074,250	02/14/2002	Laura E. Niklason	. 1579-637	5073		
23117 75	90 12/15/2003		EXAMINER			
NIXON & VANDERHYE, PC			JIANG, SHAOJIA A			
1100 N GLEBE 8TH FLOOR	ROAD		ART UNIT	PAPER NUMBER		
ARLINGTON, VA 22201-4714			1617	U		
			DATE MAILED: 12/15/2003	,		

Please find below and/or attached an Office communication concerning this application or proceeding.

. •			Application No.		Applicant(s)			
		10/074,250		NIKLASON ET AL.				
	Office Action Summary		Examiner		Art Unit			
			Shaojia A Jiang		1617			
Period fo	The MAILING DATE of this commu or Reply	inicati n appe	ears on the cover she	et with the co	orresp ndence ad	dress		
THE I - Exter after - If the - If NO - Failur - Any r	ORTENED STATUTORY PERIOD MAILING DATE OF THIS COMMUN sions of time may be available under the provision SIX (6) MONTHS from the mailing date of this conperiod for reply specified above is less than thirty period for reply is specified above, the maximum re to reply within the set or extended period for reply received by the Office later than three months dipatent term adjustment. See 37 CFR 1.704(b).	NICATION. ns of 37 CFR 1.130 nmunication. (30) days, a reply statutory period wi oly will, by statute, we	6(a). In no event, however, m within the statutory minimum (ill apply and will expire SIX (6) cause the application to becor	nay a reply be time of thirty (30) days) MONTHS from the me ABANDONED	will be considered timely the mailing date of this considered timely the mailing date of this considered this	/. mmunication.		
1)⊠	Responsive to communication(s) fi	led on <u>Septe</u>	mber 29, 2003, May	<u>12, 2003</u> .				
2a) <u></u> ☐	This action is FINAL . 2b) This action is non-final.							
3)□	Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under <i>Ex parte Quayle</i> , 1935 C.D. 11, 453 O.G. 213.							
Dispositi	on of Claims							
5)□ 6)⊠ 7)□	4) ☐ Claim(s) 1-28 is/are pending in the application. 4a) Of the above claim(s) 2-9 and 12-28 is/are withdrawn from consideration. 5) ☐ Claim(s) is/are allowed. 6) ☐ Claim(s) 1,10 and 11 is/are rejected. 7) ☐ Claim(s) is/are objected to. 8) ☐ Claim(s) are subject to restriction and/or election requirement.							
Applicati	on Papers							
9) 🗌 :	The specification is objected to by t	he Examiner						
10)	The drawing(s) filed on is/ard	e: a) acce	pted or b)☐ objected	d to by the E	xaminer.			
	Applicant may not request that any obj							
44 \ □ :	Replacement drawing sheet(s) including							
	The oath or declaration is objected under 35 U.S.C. §§ 119 and 120	to by the Exa	ammer. Note the attach	chea Office A	Action or form PT	U-152.		
12)	Acknowledgment is made of a claim All b) Some * c) None of: 1. Certified copies of the priority 2. Certified copies of the priority 3. Copies of the certified copies application from the Internative et he attached detailed Office activations as pecific reference was included of the certified copies application from the Internative et he attached detailed Office activations as pecific reference was included of CFR 1.78. The translation of the foreign lacknowledgment is made of a claim ofference was included in the first segment of the company of the comp	y documents y documents s of the priori ional Bureau on for a list of for domestic ed in the first anguage provious for domestic	have been received, have been received ty documents have b (PCT Rule 17.2(a)). If the certified copies priority under 35 U.S. sentence of the specisional application has priority under 35 U.S.	in Application peen received s.C. § 119(e) cification or its been received s.C. §§ 120 a	n No d in this National s t to a provisional n an Application s ived. and/or 121 since a	application) Data Sheet.		
	e of References Cited (PTO-892)		4) ☐ Intervi	iew Summarv (f	PTO-413) Paper No(s).		
2) 🔲 Notice	e of Draftsperson's Patent Drawing Review (nation Disclosure Statement(s) (PTO-1449)			e of Informal Pa	tent Application (PTO			

DETAILED ACTION

This application claims priority to provisional application Serial No. 60/268368.

Election/Restrictions

Applicant's election of the invention of Group I, Claims 1-23 and 28 in Paper No. 7, submitted May 12, 2003 is acknowledged.

In view of the instant claims read on <u>numerous patentably distinct agents</u> (species), the Supplemental Restriction Requirement on species election mailed July 28, 2003. In response to this Requirement, Applicant elects methotrexate as the active agent, filed in September 29, 2003 in Paper No.9, wherein Applicant also indicates that the claims readable on the elected species are claims 1, 10, and 11.

Because applicant did not distinctly and specifically point out the supposed errors in the restriction requirement, the election has been treated as an election without traverse (MPEP § 818.03(a)).

Claims 24-27 are withdrawn from further consideration pursuant to 37 CFR 1.142(b), as being drawn to a nonelected invention, there being no allowable generic or linking claim.

Claims 2-9, 12-23 and 28 are withdrawn from further consideration pursuant to 37 CFR 1.142(b), as being drawn to a non-elected species.

The claims have been examined insofar as they read on the elected specie.

Claims 1, 10, and 11 are examined on the merits herein.

Art Unit: 1617

Claim Rejections - 35 USC § 112

The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

Claim 1 is rejected under 35 U.S.C. 112, first paragraph, for scope of enablement because the specification, while being enabling for the particular agent or compound that inhibits vascular cell proliferation or particular chemotherapeutic agent disclosed in claim 11 and the specification for the claimed method herein, does not reasonably provide enablement for any compounds for inhibiting vascular cell proliferation.

These recitations, "an agent that inhibits vascular cell proliferation" and "a chemotherapeutic agent", are seen to be merely functional language.

The instant specification fails to provide information that would allow the skilled artisan to fully practice the instant invention without undue experimentation. Attention is directed to *In re Wands*, 8 USPQ2d 1400 (CAFC 1988) at 1404 where the court set forth the eight factors to consider when assessing if a disclosure would have required undue experimentation. Citing *Ex parte Forman*, 230 USPQ 546 (BdApls 1986) at 547 the court recited eight factors:

- (1) the nature of the invention; (2) the state of the prior art; (3) the relative skill of those in the art; (4) the predictability or unpredictability of the art; (5) the breadth of the claims;
- (6) the amount of direction or guidance presented; (7) the presence or absence of working examples; and (8) the quantity of experimentation necessary.

Art Unit: 1617

The nature of the invention: The instant invention pertains to a method of treating cerebral vasopasm that accompanies subarachnoid hemorrhage in a patient.

The relative skill of those in the art: The relative skill of those in the art is high.

The breadth of the claims: The instant claims are deemed very broad since the broadest claim (i.e., claim 10) reads on any compounds represented by "an agent that inhibits vascular cell proliferation" and "a chemotherapeutic agent" employed in the method herein.

The amount of direction or guidance presented:

Functional language at the point of novelty, as herein employed by Applicants, is admonished in *University of California v. Eli Lilly and Co.* 43 USPQ2d 1398 (CAFC, 1997) at 1406: stating this usage does "little more than outline goal appellants hope the recited invention achieves and the problems the invention will hopefully ameliorate". The CAFC further clearly states that "[A] written description of an invention involving a chemical genus, like a description of a chemical species, requires a precise definition, such as by structure, formula, [or] chemical name, of the claimed subject matter sufficient to distinguish it from other materials" at 1405(emphasis added), and that "It does not define any structural features commonly possessed by members of the genus that distinguish from others. One skilled in the art therefore cannot, as one can do with a fully described genus, visualize or recognize the identity of the members of the genus. A definition by function, as we have previously indicated, does not suffice to define the genus.." at 1406 (emphases added).

Art Unit: 1617

In the instant case, "an agent that inhibits vascular cell proliferation" and "a chemotherapeutic agent", recited in the instant claims are purely functional distinction. Hence, these functional recitations read on any compounds that might have the recited functions. However, the specification merely provides one particular compound for each kind of functional compounds in the specification.

Thus, Applicants functional language at the points of novelty fails to meet the requirements set forth under 35 U.S.C. 112, first paragraph. Claims employing functional language at the exact point of novelty, such as Applicants', neither provide those elements required to practice the inventions, nor "inform the public during the life of the patent of the limited of monopoly asserted" (*General Electric Company v. Wabash Appliance Corporation et al.* 37 USPQ at 468 (US Supreme Court 1938)).

The predictability or unpredictability: the instant claimed invention is highly unpredictable as discussed below:

It is noted that the pharmaceutical art is <u>unpredictable</u>, requiring each embodiment to be individually assessed for physiological activity. *In re Fisher*, 427 F.2d 833, 166 USPQ 18 (CCPA 1970) indicates that the more unpredictable an area is, the more specific enablement is necessary in order to satisfy the statute. In the instant case, the instant claimed invention is highly <u>unpredictable</u> since one skilled in the art cannot fully described genus, visualize or recognize the <u>identity</u> of the members of the genus, by structure, formula, or chemical name, of the claimed subject matter, as discussed above in *University of California v. Eli Lilly and Co.* Hence, in the absence of fully recognizing the identity of the members genus herein, one of skill in the art would

Art Unit: 1617

be <u>unable</u> to fully predict possible physiological activities of any compounds having claimed functional properties in the pharmaceutical compositions herein.

Moreover, one of skill in the art would recognize that it is highly unpredictable in regard to therapeutic effects, side effects, and especially serious toxicity that may be generated by drug-drug interactions when and/or after administering to a host (e.g., a human) any compounds represented by "an agent that inhibits vascular cell proliferation" and "a chemotherapeutic agent", which may encompass more than a thousand compounds. See text book "Goodman & Gilman's The Pharmacological Basis of Therapeutics" regarding possible drug-drug interactions (9th ed, 1996) page 51 in particular. This book teaches that "The frequency of significant beneficial or adverse drug interactions is unknown" (see the bottom of the left column of page 51) and that "Recognition of beneficial effects and recognition of and prevention of adverse drug interactions require a thorough knowledge of the intended and possible effects of drugs that are prescribed" and that "The most important adverse drug-drug interactions occur with drugs that have serious toxicity and a low therapeutic index, such that relatively small changes in drug level can have significant adverse consequences" (see the right column of page 51) (emphases added). In the instant case, in the absence of fully recognizing the identity of the members genus herein, one of skill in the art would not be able to fully predict possible adverse drug-drug interactions occurring with many combinations of any compounds having claimed functional properties in the pharmaceutical compositions herein to be administered to a host. Thus, the teachings of the book clearly support that the instant claimed invention is highly unpredictable.

Art Unit: 1617

The presence or absence of working examples and the quantity of experimentation necessary:

As discussed above, only one particular compound for each kind of functional compounds employed in the composition herein is disclosed in the specification.

Moreover, it is noted that the specification fails to provide working examples, i.e., testing results or data to demonstrate that any chemotherapeutic agent or methotrexate to be administered to a host, i.e., in vitro or vivo, in treating for cerebral vasopasm in a patient.

Thus, the specification fails to provide sufficient support of the broad use of any compounds represented by "an agent that inhibits vascular cell proliferation" and "a chemotherapeutic agent" recited in the instant claims. As a result, necessitating one of skill to perform an exhaustive search for the embodiments of <u>any</u> compounds having those functions recited in the instant claims suitable to practice the claimed invention.

Genentech, 108 F.3d at 1366, states that "a patent is not a hunting license. It is not a reward for search, but compensation for its successful conclusion" and "[p]atent protection is granted in return for an enabling disclosure of an invention, not for vague intimations of general ideas that may or may not be workable".

Therefore, in view of the <u>Wands</u> factors, the case <u>University</u> of <u>California v. Eli</u>

Lilly and Co. (CAFC, 1997) and <u>In re Fisher</u> (CCPA 1970) discussed above, to practice the claimed invention herein, a person of skill in the art would have to engage in <u>undue</u>

<u>experimentation</u> to test all compounds encompassed in the instant claims and their

Art Unit: 1617

combinations employed in the claimed compositions to be administered to a host, with no assurance of success.

Claim Rejections - 35 USC § 112

The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

Claims 1 and 10-11 are rejected under 35 U.S.C. 112, first paragraph, for scope of enablement because the specification, while being enabling for a method of treating cerebral vasopasm that accompanies subarachnoid hemorrhage in a patient disclosed in the specification employing the instant compounds herein, does not reasonably provide-enablement-for-the-**prevention** or **preventing** cerebral vasopasm that-accompanies subarachnoid hemorrhage in a patient.

The instant claims are drawn to the methods of <u>preventing</u> cerebral vasopasm that accompanies subarachnoid hemorrhage in a patient. The instant specification <u>fails</u> to provide information that would allow the skilled artisan to practice the instant invention. Attention is directed to *In re Wands*, 8 USPQ2d 1400 (CAFC 1988) at 1404 where the court set forth the eight factors to consider when assessing if a disclosure would have required undue experimentation. Citing *Ex parte Forman*, 230 USPQ 546 (BdApls 1986) at 547 the court recited eight factors:

(1) the nature of the invention; (2) the state of the prior art; (3) the relative skill of those in the art; (4) the predictability or unpredictability of the art; (5) the breadth of the claims;

Art Unit: 1617

(6) the amount of direction or guidance presented; (7) the presence or absence of working examples; and (8) the quantity of experimentation necessary.

Nature of the invention: The instant invention pertains to the method of preventing cerebral vasopasm that accompanies subarachnoid hemorrhage in a patient i.e., a human or animal.

The state of the prior art: The skilled artisan would view that the treatment to prevent cerebral vasopasm that accompanies subarachnoid hemorrhage in a human or animal totally, absolutely, or permanently, is highly unlikely, not even occurring at the first time.

The relative skill of those in the art: The relative skill of those in the art is high.

The predictability or lack thereof in the art: The skilled artisan would view that the treatment to prevent cerebral vasopasm that accompanies subarachnoid hemorrhage in a human or animal totally, absolutely, or permanently is highly unpredictable, and not even occur at the first time is highly unpredictable.

The amount of direction or guidance presented and the presence or absence of working examples: In the instant case, <u>no</u> working examples are presented in the specification as filed originally showing how to prevent cerebral vasopasm that accompanies subarachnoid hemorrhage in a human or animal totally, absolutely, or permanently, not even occurring at the first time. As discussed above, preventing prevent cerebral vasopasm that accompanies subarachnoid hemorrhage in a human or animal totally, absolutely, or permanently is highly unpredictable and unlikely, and not even occur at the first time is highly unpredictable.

Art Unit: 1617

Genentech, 108 F.3d at 1366, states that "a patent is not a hunting license. It is not a reward for search, but compensation for its successful conclusion" and "[p]atent protection is granted in return for an enabling disclosure of an invention, not for vague intimations of general ideas that may or may not be workable".

Therefore, in view of the <u>Wands</u> factors, e.g., the amount of direction or guidance provided, absence of working examples, and the predictability of the art discussed above, to practice the claimed invention herein, a person of skill in the art would have to engage in <u>undue experimentation</u> to test all compounds encompassed in the instant claims to be administered to a host employed in the claimed methods of preventing cerebral vasopasm that accompanies subarachnoid hemorrhage, with no assurance of success.

Claim Rejections - 35 USC § 102

The following is a quotation of the appropriate paragraphs of 35 U.S.C. 102 that form the basis for the rejections under this section made in this Office action:

A person shall be entitled to a patent unless -

(b) the invention was patented or described in a printed publication in this or a foreign country or in public use or on sale in this country, more than one year prior to the date of application for patent in the United States.

Claims 1 and 10-11 are rejected under 35 U.S.C. 102(b) as being anticipated by Black (5,527,778, PTO-892).

Black discloses that well-known neuropharmaceutical agents such as chemotherapeutic agents, in particular, methotrexate (see col.4 line 57 to col.5 line 10) to be administered to a patient are useful in methods of treating abnormal brain tissue including subarachnoid hemorrhage, head injury (head trauma) and cerebral ischemia,

Art Unit: 1617

and opening abnormal brain tissue capillaries in a patient, i.e., a mammal (see abstract, col.4 lines 1-9).

Thus, the disclosure of Black anticipates claims 1 and 10-11.

In view of the rejections to the pending claims set forth above, no claims are allowed.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Examiner Jiang, whose telephone number is (703) 305-1008. The examiner can normally be reached on Monday-Friday from 9:00 to 5:00.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Sreenivasan Padmanabhan, Ph.D., can be reached on (703) 305-1877.

The fax phone number for the organization where this application or proceeding is assigned is (703) 308-4556.

Any inquiry of a general nature or relating to the status of this application or proceeding should be directed to the receptionist whose telephone number is (703) 305-

1235.

S. Anna Jiang, Ph.D.

Patent Examiner, AU 1617

December 8, 2003